10

15

20

25

30

Metered dose inhaler for salmeterol xinafoate

The invention provides a container for a metered dose inhaler (MDI) for use in dispensing a quantity of a pharmaceutical formulation, especially salmeterol xinafoate formulation, which may be used in the treatment of respiratory disorders.

4-hydroxy- α^1 -[[[6-(4-phenylbutoxy)hexyl]amino]methyl]-1,3-benzenedimethanol was described as one of a wide range of bronchodilators in GB-A-2140800. This compound is also known by the generic name of salmeterol, the 1-hydroxy-2-naphthoate (xinafoate) salt of which has become widely known as a highly effective treatment of inflammatory diseases, such as asthma and chronic obstructive pulmonary disease (COPD).

Containers for aerosol formulations commonly comprise a vial body (canister) coupled to a valve. The valve comprises a valve stem through which the formulations are dispensed. Generally the valve includes a rubber valve seal intended to allow reciprocal movement of the valve stem which prevents leakage of propellant from the container. Metered dose inhalers comprise a valve which is designed to deliver a metered amount of an aerosol formulation, to the recipient, per actuation. Such a metering valve generally comprises a metering chamber which is of a set volume which aims to administer per actuation an accurate, predetermined dose.

Metering valves incorporate gaskets (also known as seals) to prevent leakage of propellant through the valve. The gasket may comprise suitable elastomeric material such as for example low density polyethylene, chlorobutyl, black and white butadiene-acrylonitrile rubbers, butyl rubber and neoprene.

Valves for use in MDIs are available from manufacturers well known in the aerosol industry, for example, from Valois, France (eg. DF10, DF30, DF60), Bespak plc, United Kingdom (eg. BK300, BK356, BK357) and 3M-Neotechnic Limited, United Kingdom (eg. SpraymiserTM). The metering valves are used in association with commercially available canisters, for example metal canisters, such as aluminium canisters, suitable for delivering pharmaceutical aerosol formulations.

MDIs incorporating valve gaskets as described above perform adequately with chlorofluorocarbon propellants such as propellant 11 (CCl $_3$ F), propellant 114 (CF $_2$ ClCF $_2$ Cl) and propellant 12 (CCl $_2$ F $_2$) or mixtures thereof. However these propellants are now believed to provoke the degradation of stratospheric ozone and there is thus a need to provide aerosol formulations for medicaments which employ so called "ozone-friendly" propellants.

A class of propellants which are believed to have minimal ozone-depleting effects in

10

15

20

25

comparison to conventional chlorofluorocarbon propellants comprises hydrofluoroalkanes (HFA's) especially 1,1,1,2-tetrafluoroethane (HFA134a), 1,1,1,2,3,3,3-heptafluoro-n-propane (HFA 227) and mixtures thereof. However there have been problems associated with stabilising the pharmaceutical aerosol formulations prepared using the new class of propellants.

Pharmaceutical aerosol formulations may comprise a solution or a suspension. Some solution formulations suffer the disadvantage that the drug substance contained therein is more susceptible to degradation. Furthermore there are issues with control of size of the droplets which influences the therapeutic profile. For this reason, suspensions are generally preferred.

Suspension aerosol formulations generally comprise a particulate medicament, one or more liquid propellants, optionally with a co-propellant, and optionally an adjuvant such as a solvent or a surfactant. The aerosol formulation is under pressure in the canister.

The efficiency of an aerosol device, such as an MDI, is a function of the dose deposited at the appropriate site in the lungs. Deposition is affected by several factors, of which one of the most important is the aerodynamic particle size. Solid particles and/or droplets in an aerosol formulation can be characterised by their mass median aerodynamic diameter (MMAD, the diameter around which the mass aerodynamic diameters are distributed equally).

In suspension formulations, particle size in principle is controlled during manufacture by the size to which the solid medicament is reduced, usually by micronisation. However, if the suspended drug has the slightest solubility in propellant, a process known as Ostwald Ripening can lead to particle size growth. Also, particles may have the tendency to aggregate, or adhere to parts of the MDI eg. canister or valve. Furthermore the drug may have the tendency to be absorbed into the rubber components of the valve, especially when stored for a prolonged period. In particular fine particles may be preferentially absorbed. The effect of Ostwald Ripening and especially of drug deposition may be particularly severe for potent drugs (including salmeterol xinafoate) which need to be formulated in low doses.

The problems mentioned above have been addressed by the addition of one or more of adjuvants such as alcohols, alkanes, dimethyl ether, surfactants (including fluorinated and non-fluorinated surfactants, carboxylic acids, polyethoxylates etc) and even conventional chlorofluorocarbon propellants in small amounts intended to minimise potential ozone damage as disclosed in, for example, EP0372777, WO91/04011, WO91/11173, WO91/11495 and WO91/14422.

10

15

20

25

30

35

Excipient free formulations of salmeterol xinafoate are described in WO93/11743.

Furthermore, WO96/32345, WO96/32151, WO96/32150 and WO96/32099 disclose aerosol canisters coated with one or more fluorocarbon polymers optionally in combination with one or more non-fluorocarbon polymers.

It is essential that the prescribed dose of aerosol medication delivered from MDIs to the patient consistently meet the specifications claimed by the manufacturer and comply with the requirements of the FDA and other regulatory authorities. That is, every dose delivered from the can must be the same within close tolerances. Therefore it is important that the formulation be substantially homogeneous throughout the administered dose throughout the life of the product. It is also important that the concentration of the suspension does not change significantly when stored for a prolonged period.

To obtain regulatory approval, pharmaceutical aerosol formulation products must meet strict specifications. One parameter for which a specification is usually set is the fine particle mass (FPM). This is a means of evaluating the amount of drug substance which has the potential to reach the inner lungs, i.e. the small bronchioles and alveoli, based on the amount of drug particles with a diameter within a certain range, usually less than 5 microns.

The FPM of an actuation from an MDI can be calculated based on the sum of the amount of drug substance deposited on stages 3, 4 and 5 of an Andersen Cascade Impaction stack as determined by standard HPLC analysis.

It is important that the FPM of the pharmaceutical aerosol formulation for all the doses dispensed from the MDI are within the specification set, even after the MDI has been stored for a prolonged period.

Whilst not wishing to be bound by any theories it is hypothesised by the inventors that the concentration of drug in the suspension and thus the dose dispensed may, in many cases (especially in particulate salmeterol xinafoate and HFA formulations), decrease over time with the adsorption of drug into the rubber components of the valve. This may be observed as a decrease in the Total Drug Content (TDC) of the can. This process may be accelerated by the ingression of water into the formulation.

This hypothesis has been supported by studies employing salmeterol xinafoate HFA 134a aerosol formulations in conventional MDI's stored at 40°C and 75% relative humidity and 40°C and 20% relative humidity as shown in Table 1.

Furthermore evidence indicates that the FPM and mean dose of some particulate aerosol formulations, for example, salmeterol xinafoate HFA 134a formulations

10

15

20

decreases over time with the ingression of water into the formulation and/or deposition and/or absorption resulting in impaired performance of the MDI.

The effect on FPM of salmeterol xinafoate HFA 134a aerosol formulations in conventional MDIs stored at 40°C and 75% relative humidity is shown in Table 2 and Table 4. Table 3 and Table 5 show a noticeable decrease over time in the mean dose delivered from a conventional MDI when stored at 40°C and 75% relative humidity.

Deposition of drug particles on other valve components, particularly the metering chamber may also contribute to the formulation stability problems observed such as inconsistencies in the doses dispensed, which becomes particularly acute over increasing numbers of actuations.

The problem with deposition is particularly exacerbated when excipient-free aerosol formulations are used based on the propellants 1,1,1,2-tetrafluoroethane (HFA 134a) and 1,1,2,3,3,3-heptafluoro-n-propane (HFA 227) and is thought to increase with length of storage of the aerosol, particularly when stored at high temperature and/or high humidity.

Surprisingly the inventors have found that the suspension concentration, dose and FPM of formulations of particulate medicament suspended in a HFA propellant e.g. salmeterol xinafoate in suspension in a HFA propellant, obtained from an MDI may be stabilised by reducing the deposition on the valve component(s) and reducing the drug absorption into rubber components and/or effectively controlling the ingression of water into the formulation during storage and use by employing particular valve materials.

Thus the invention provides a container comprising a canister sealed with a metering valve having a metering chamber, which contains a pharmaceutical aerosol formulation consisting essentially of

- 25 (A) particulate salmeterol xinafoate optionally in combination with another drug useful in inhalation therapy, suspended in
 - (B) a liquefied propellant gas comprising 1,1,1,2,3,3,3-heptafluoro-n-propane, 1,1,1,2-tetrafluoroethane or a mixture thereof;

wherein the formulation is substantially free of surfactant and components having polarity higher than the liquefied propellant gas;

said valve characterised in that it contains one or more sealing gaskets substantially constructed from of a polymer of ethylene propylene diene monomer (EPDM) and the metering chamber surface presents a substantially fluorinated surface to the formulation.

10

15

25

30

35

Preferably the formulation will consist of particulate salmeterol xinafoate optionally in combination with another drug useful in inhalation therapy, suspended in 1,1,1,2,3,3,3-heptafluoro-n-propane, 1,1,1,2-tetrafluoroethane or a mixture thereof.

More preferably the liquefied propellant gas is 1,1,1,2-tetrafluoroethane.

We have found that the trends observed, especially for salmeterol xinafoate, wherein the dose delivered and the FPM is reduced after storage, especially at elevated temperatures and under high humidity conditions, can be ameloriated by use of one or more gaskets constructed substantially from a polymer of EPDM. However the absolute values of the dose delivered and the FPM are not significantly increased. Therefore to ensure the patient gets the correct dose each time the device is actuated the formulation must contain an excess of drug substance, sometimes called "overage", to compensate for the loss. Advantageously when said gaskets are used in conjunction with a metering chamber which presents a substantially fluorinated surface to the formulation, the absolute dose of medicament available to the patient is raised whilst simultaneously maintaining or improving the stabilisation in the dose delivered and FPM. This provides benefits to the patient who receives the full dose claimed to be available on the label of the medicament and is more likely to satisfy the rigorous standards of the FDA and other regulatory authorities. Furthermore there are economic advantages since wastage of product is reduced.

Furthermore water is repelled from the fluorinated surface of the metering chamber which may further reduce the water ingression into the formulation over time, thereby reducing the undesirable effects thereof.

A particular aspect of the invention is a container as described above wherein the valve is sealed to the canister by means of a can/neck sealing gasket (3) which is substantially constructed from a polymer of EPDM.

Especially preferred is a container as described above wherein the metering valve comprises a metering chamber (4) defined by walls and an upper (12) and a lower (9) sealing gasket through which pass a valve stem (7 and 8) characterised in that said two sealing gaskets are substantially constructed from a polymer of EPDM and the metering chamber surface presents a substantially fluorinated surface to a formulation containable therein.

Also especially preferred is a container as described above wherein the valve is sealed to the canister by means of a can sealing gasket (3) which is substantially constructed from EPDM polymer and wherein the lower (9) sealing gasket is substantially constructed from EPDM polymer.

10

15

20

25

30

Most preferably all the sealing gaskets in the said metering valve are substantially constructed from EPDM polymer.

In the foregoing, the expressions "polymer of EPDM" and "EPDM polymer" are used interchangeably.

Sealing gasket when used in this specification will be understood to mean a neck/canister gasket and/or lower sealing gasket and/or upper sealing gasket.

Figure 1 shows part of a cross-section view of an MDI, with the valve pointing downward. The gaskets are represented by: 3 the can/neck seal, 9 the lower metering chamber seal and 12 the upper metering chamber seal. The metering chamber is identified as 4 and the stem is identified as 7 and 8.

Figure 2 shows an alternative cross-section of an MDI valve.

EPDM polymer when used as a gasket material in valves for use with aerosol formulations of particulate medicament in a HFA propellant appears to reduce deposition of drug particles on said gaskets in comparison to those gaskets prepared from traditional materials.

Furthermore EPDM polymer properties have been found to be superior to those materials traditionally used with respect to the absorption of drug into rubber.

In addition it seems that EDPM polymer may also have superior properties with respect to the control of water ingression into the pharmaceutical aerosol formulation containing hydrofluorocarbons. This is illustrated in Table 2 which shows that salmeterol xinafoate HFA 134a formulations in MDIs with gaskets of EPDM polymer have a stable FPM and dose delivered at the beginning of use even when stored at 40°C and relative humidity 75% for up to 6 months.

Table 3 and Table 5 give mean dose data and range of dose data for beginning of use which further supports the improved stability of formulations illustrated by salmeterol xinafoate HFA 134a formulations wherein the valve gaskets are composed of EPDM polymer.

In addition it seems that the life span of the gaskets of EPDM polymer is longer than that of traditional gaskets as the material is more stable to the hydrofluorocarbon containing formulations and more resistant to degradation and/or distortion. Therefore the advantages of the EPDM polymer are enjoyed throughout the life of the product without the function of the device being impaired.

EPDM polymer is available from a variety of suppliers including West and Parker Seals (USA).

A gasket substantially constructed from a polymer of EPDM when used in this

10

25

30

specification will be understood to mean a gasket composed of greater than 90% of EPDM polymer, particularly greater than 95% of EPDM polymer, especially greater than 99% of EPDM polymer.

The invention also relates to a container as described above wherein the metering chamber presents a substantially fluorinated surface to the formulation. This advantageously reduces drug deposition on the metering chamber when used in conjunction with aerosol HFA formulations such as salmeterol xinafoate HFA formulations, compared with valves conventionally available.

The said metering chamber may be constructed from of any material with suitable characteristics such as any conventionally used plastics material such as nylon, polybutylene terephthalate PBT (polyester), acetal (polyoxymethylene) and tetrabutyrene terephthalate (TBT) etc, or metallic material which is compatible for use with the formulation, for example, stainless steel or aluminium. One example of a metal valve is the 3M-Neotechnic valve.

The metering chamber (especially when composed of a plastics material) is preferably surface treated so as to present a substantially fluorinated surface to the formulation. Preferably surface treatment will comprise a process of plasma coating with highly fluorinated small molecules such as: C₁₋₁₀perfluoroalkanes including perfluorocycloalkanes; C₂₋₁₀perfluoroalkenes; fluoroalkanes including fluorocycloalkanes or fluoroalkenes wherein a high proportion of the hydrogens have been replaced by fluorines or mixtures thereof. Furthermore the fluorinated molecules or mixtures thereof

Especially preferred small molecules include C₁₋₁₀perfluoroalkanes.

The plasma coating may comprise a fluorinated polymer laid down on the surface of the valve component, preferably the metering chamber, by polymerisation or direct modification of the material surface by interchange of hydrogen ions in the material with fluorine ions. The coating process typically takes place in a vacuum at ambient temperature. The components to be coated are placed inside a chamber which is evacuated. The fluorine monomer or fluorine source is introduced into the chamber at a controlled rate. The plasma is ignited within the chamber and maintained for a given time at a chosen power setting. At the end of the treatment the plasma is extinguished, the chamber flushed and the products retrieved. In the polymerisation process, a thin layer of plasma polymer will be bonded to the surface of the valve component, preferably a metering chamber, or any other surface of the valve to be coated.

may optionally be used in combination with one or more non-fluorocarbon compounds.

For plasma polymerization typically temperatures in the range of about 20°C to about

10

15

20

25

100°C may be employed.

The surface of the component especially the metering chamber may require activating in order to facilitate more effective coating of the surface by improving the adhesion of the coating to the surface.

Preferably the components to be plasma coated will be pre-treated to remove any surface contamination and/or to activate the surface. This may be achieved by, for example, treatment of the components with an etching gas such as oxygen or argon. In the process radicals react with the plastic or metal substrate e.g the component is exposed to a low pressure oxygen plasma environment which creates polar groups on the components surface which are more conducive to bonding with the plasma coating to be applied.

Alternatively the metering chamber (especially when composed of a plastics material, for example, those described above) may be surface treated with a siloxane such as dimethyl siloxane using a similar process as that described above for fluoroplasma coating.

Alternatively the metering chamber presents a substantially fluorinated surface to the formulation by virtue of being composed of a suitable substantially fluorinated material. Suitable fluorinated materials include fluorinated polymer/copolymer or mixtures thereof or a mixture of the fluorinated polymer in combination with non-fluorinated polymers conventionally used in the manufacture of valves, such as acetal, polyester (PBT). Examples of suitable fluorinated polymers include polytetrafluoroethylene (PTFE), ethylenetetrafluoroethylene (ETFE), polyvinyldienefluoride (PVDF), perfluoroalkoxyalkane (PFA), polyvinylfluoride (PVF), polychlorotrifluoroethylene (PCTFE), fluorinated ethylenepropylene (FEP) etc. Suitable copolymers include copolymers of tetrafluoroethylene (TFE) with PFA, TFE with hexafluoropropylene (HFP) (available as FEP 6107 and FEP 100 from DYNEON), VDF with HFP (commercially available as Viton A), TFE with perfluoro(propyl vinyl ether) (available as PFA 6515N from DYNEON), a blend of TFE, hexafluoropropylene and vinylidene fluoride (available commercially as THV 200G from DYNEON), etc.

It should be noted, however, that any conventionally available polymer, copolymer or mixture thereof which comprises a fluorinated polymer and which can be used to make the valve for use in an inhaler according to the invention may be suitable. Examples of mixtures of polymers and/or copolymers comprise, for example, up to 80% by weight fluorinated polymer, optionally up to 40% by weight fluorinated polymer, optionally up to 5% by weight of fluorinated 20% by weight fluorinated polymer or optionally up to 5% by weight of fluorinated

10

15

20

25

30

35

polymer. Preferably, fluorinated polymers selected from PTFE, PVF and PCTFE are used as mixtures with non-fluorinated polymers. For example a suitable material is HOSTAFORM X329TM (Hoechst) which is a 5% PTFE/Acetal blend, HOSTAFORM C9021TF which is a 20% PTFE/Acetal blend, PTFE/PBT blends (for example, LNP WL4040), PTFE/PBT/silicone blends (for example, LNP WL4540).

The fluorinated polymers and mixtures thereof used in the invention can be moulded in any conventional manner, for example, by injection moulding, plastic moulding etc.

Alternatively metering chambers (especially when composed of a metallic material such as aluminium or stainless steel) can be coated by conventional techniques using fluorocarbon polymers which include fluorocarbon polymers which are made of multiples of one or more of the following monomeric units: tetrafluoroethylene (PTFE), fluorinated ethylene propylene (FEP), perfluoroalkoxyalkane (PFA), ethylene tetrafluoroethylene (ETFE), vinyldienefluoride (PVDF), and chlorinated ethylene tetrafluoroethylene. Fluorinated polymers, which have a relatively high ratio of fluorine to carbon, such as perfluorocarbon polymers, e.g. PTFE, PFA or FEP may be preferable, especially polymers selected from PTFE and FEP.

The metering chamber may be treated, so as to present a substantially fluorinated surface to the formulation, for example, by coating with a fluorinated polymer which is optionally blended with non-fluorinated polymers such as polyamides, polyimides, polyamideimides, polyethersulfones, polyphenylene sulfides, and amine-formaldehyde thermosetting resins. These added polymers often improve adhesion of the polymer coating to the substrate. Preferred polymer blends are PTFE/FEP/polyamideimide, PTFE/ polyethersulphone (PES) and FEP-benzoguanamine. The most preferred polymer coating is a blend of PTFE and PES. A coating of pure FEP is also of considerable interest.

A technique for applying such coatings to, for example, a metal, such as aluminium or stainless steel, is where the metal is precoated as coil stock and cured before being stamped or drawn into the can shape. This method is well suited to high volume production for two reasons. First, the art of coating coil stock is well developed and several manufacturers can custom coat metal coil stock to high standards of uniformity and in a wide range of thicknesses. Second, the precoated stock can be stamped or drawn at high speeds and precision by essentially the same methods used to draw or stamp uncoated stock.

Other techniques for coating techniques includes electrostatic dry powder coating or by spraying preformed MDI components with formulations of the coating fluorinated

10

15

polymer/polymer blend and then curing. The preformed MDI components may also be dipped in the fluorocarbon polymer/polymer blend coating formulation and cured, thus becoming coated on the inside and out. The fluorocarbon polymer/polymer blend formulation may also be poured inside the MDI components then drained out leaving the insides with the polymer coat.

The appropriate curing temperature is dependent on the polymer blend chosen for the coating and the coating method employed.

However, for coil coating and spray coating temperatures in excess of the melting point of the polymer are typically required, for example, about 50°C above the melting point for up to about 20 minutes such as about 5 to 10 minutes e.g. about 8 minutes or as required. For the above named preferred and particularly preferred polymer blends curing temperatures in the range of about 300°C to about 400°C, e.g. about 350°C to 380°C are suitable.

Where the components are coated and then cured the substrate components may be prepared from strengthened materials to ensure they withstand the process.

Thus an aspect of the invention is a process for preparing a container, as described above, wherein the surface treatment of the metering chamber comprises a process for applying a coating of a fluorocarbon polymer optionally in combination with a non-fluorocarbon polymer.

Conversely alternative polymer coatings may be used on the components which may be dipped or bath immersed into a treatment tank containing a solution of polymeric compound. Usually the components are immersed in the solution at room temperature for at least one hour, for example, 12 hours, thus being treated both internally and externally.

The treated components are preferably washed with solvent and dried at an elevated temperature for example 50-100°C optionally under vacuum.

Examples of suitable coating materials include of fluoropolyethers having functionalised ends groups with a general formula $R_FO(C_3F_6O)_m(CFX)_n$ -CFX-Y- Z_p as described in USP 4, 746, 550 (incorporated herein by reference) including perfluoropolyethers having functional groups capable of anchoring the coating to the substrate such as carboxyl, ester, amide, hydroxyl, isocyanate, epoxy, silane, for example, -CONR²R³ wherein R² and R³ may be independently selected from amongst other things hydrogen, or a silyl ether (e.g. $SiR_1(OR)_{3-1}$ or a fluoropolyether having hydroxylic functionality of the type -CF₂CH₂OH, -CF₂CFXCH₂OH (wherein X is Cl or F) or -CF(CF₃)CH₂OH as described in USP 6, 071, 564 (incorporated herein by reference); phosphoric diesters of formula

30

10

20

25

30

35

[XCF₂CF₂O(CFXCF₂O)_xCFXCH₂O]₂PO(OM) as described in USP 3, 492, 374 (incorporated herein by reference) or phosphoric monoester of formula [R_f-O-CFY-L-O]_mP=O(O'Z⁺)_{3-m} as described in EP 0 687 533 (incorporated herein by reference) wherein L is a divalent organic group; m = 1; Y is -F or $-CF_3$; Z^+ is selected from H⁺, M⁺ where M is an alkali metal; N(R)₄⁺ where the R groups independently represent H or C₁₋₆alkyl; R_f is a polyperfluoroalkyleneoxide chain.

The fluoropolyethers described above may be used in combination with monofunctional fluoropolyethers having $-CH_2OH$ terminals directly linked to a perfluoroalkyl group $-CF_2$, $-CF_2CFX$ (wherein X is Cl or F) or $CF(CF_3)$ optionally through a bridging group $(CH_2CH_2)_q$ wherein q represents an integer from 1 to 6.

Other suitable coating materials also include polymeric compounds that are silane derivatives of perfluoropolyoxyalkanes with a molecular weight in the range 1600-1750 and those of the general formula:

15
$$R^1 - (CH_2)_v - CF_2O - (C_2F_4O)_x - (CF_2O)_v CF_2 - (CH_2)_w - R^1$$
 (I)

wherein R¹ comprises:

- $(OCH_2-CH_2)_z$ - $OPO(OH)_2$, wherein x, y and z are such that the molecular weight of the compound is 900-2100 and v and w independently represent 1 or 2.

The invention also relates to a container as described above wherein the valve stem presents a substantially fluorinated surface to the formulation.

Stems to be plasma coated may optionally be pretreated to remove surface contamination and/or activate the surface.

Alternatively stems may be coated by conventional techniques using fluorocarbon polymers optionally in combination with non-fluorocarbon polymer wherein the said coating requires curing after application as described above.

Additionally stems may be coated by processes using fluorocarbon polymers that require drying at temperatures between 50-100°C as described above for metering chambers.

Alternatively the stem presents a substantially fluorinated surface to the formulation by virtue of being composed of a suitable fluorinated material.

Analogous processes and materials described above for metering chambers are suitable for the preparation of valve stems according to the invention.

Preferably the substantially fluorinated surface will result from surface treatment of the stem. Most preferably the surface treatment will comprise a process of plasma coating with highly fluorinated small molecules such as: C₁₋₁₀perfluoroalkanes including

20

25

fluorocycloalkanes; C_{2-10} perfluoroalkenes; fluoroalkanes including fluorocycloalkanes or fluoroalkenes wherein a high proportion of the hydrogens have been replaced by fluorines or mixtures thereof as described above.

Preferably the container according to the invention is a canister composed of aluminium.

Preferably the canister also presents a substantially fluorinated surface to the formulation.

Preferably the canister is surface treated so as to present a substantially fluorinated surface to the formulation contained therein.

More preferably the canister is surface treated by coating with a fluorocarbon polymer optionally in combination with a non-fluorocarbon polymer, for example, using materials mentioned above. Fluorocarbon polymers selected from FEP and PTFE are particularly preferred for the surface treatment of canisters. FEP is especially preferred. A polymer blend of PTFE and PES is also especially preferred.

The surface treatment of the canister may be performed by methods analogous to those described above for valve components.

Preferably salmeterol xinafoate is the only medicament. However medicaments which may be administered in aerosol formulations according to the invention in combination with salmeterol xinafoate include any drug useful in inhalation therapy e.g; anti-allergics, e.g. cromoglycate (e.g. as the sodium salt), ketotifen or nedocromil (e.g. as sodium salt); anti-inflammatory steroids, e.g. beclomethasone (e.g. as dipropionate), fluticasone (e.g. as propionate), flunisolide, budesonide, rofleponide, mometasone (e.g. as furoate), ciclesonide, triamcinolone acetonide; anticholinergics, e.g. ipratropium (e.g. as bromide), tiotropium, atropine or oxitropium and salts thereof. It will be clear to a person skilled in the art that, where appropriate, the medicaments may be used in the form of salts, (e.g. as alkali metal or amine salts or as acid addition salts) or as esters (e.g. lower alkyl esters) or as solvates (e.g. hydrates) to optimise the activity and/or stability of the medicament and/or to minimise the solubility of the medicament in the propellant.

30 Specific medicaments of interest for use in combination with salmeterol xinafoate include fluticasone propionate or ipratropium bromide.

The container and valve described herein may also be suitable for containing medicaments besides salmeterol xinafoate which present similar formulation difficulties e.g. because of their susceptibility to water ingress, drug deposition, and other drug losses. Generally these difficulties are especially severe for potent medicaments which

15

20

25

30

are administered at low doses. Examples include salmeterol and salts thereof, fluticasone propionate, formoterol and salts thereof. Other example medicaments include beclomethasone dipropionate, budesonide, sodium cromoglycate, albuterol and salts thereof and combinations thereof.

Medicament may be used in the form of racemate or in the form of a pure isomer e.g. R-salmeterol or S-salmeterol.

The particle size of the particulate (e.g. micronised) medicament should be such as to permit inhalation of substantially all of the medicament into the lungs upon administration of the aerosol formulation and will thus be less than 100 microns, desirably less than 20 microns, and preferably in the range 1-10 microns, e.g. 1-5 microns.

The concentration of medicament in the formulation will generally be 0.01-10% such as 0.01-2%, particularly 0.01-1%, especially 0.03-0.25% w/w. When salmeterol xinafoate is the only medicament its concentration in the formulation will generally be 0.03-0.15% w/w.

It is desirable that the formulations of the invention contain no components which may provoke the degradation of stratospheric ozone. In particular it is desirable that the formulations are substantially free of chlorofluorocarbons such as CCl₃F, CCl₂F₂ and CF₃CCl₃. It is desirable that the formulations of the invention are substantially free of any volatile adjuvant such as a saturated hydrocarbon, for example, propane, n-butane, isobutane, pentane and isopentane or a dialkyl ether, for example, dimethyl ether.

It is desirable that the formulations of the invention are substantially free of surfactant. "Substantially free" will generally be understood to mean containing less than 0.01% w/w especially less than 0.0001% based on weight of medicament.

It is desirable that the formulations of the invention are substantially free of any polar adjuvants e.g. C₂₋₆aliphatic alcohols and polyols such as ethanol, isopropanol propylene glycol, glycerol and mixtures thereof. "Substantially free" will generally be understood to mean containing less than 0.01% especially less than 0.0001% based on weight of formulation. Polarity may be determined, for example, by the method described in European Patent Application Publication No. 0327777.

Thus in one aspect the invention provides a container which contains a pharmaceutical aerosol formulation comprising a particulate medicament and a liquefied propellant gas of 1,1,1,2,3,3,3-heptafluoro-n-propane, 1,1,1,2-tetrafluoroethane or mixtures thereof. Preferably the pharmaceutical aerosol formulation will consist of or consist essentially of

20

25

30

a particulate medicament and a liquefied propellant gas of 1,1,1,2,3,3,3-heptafluoro-n-propane, 1,1,1,2-tetrafluoroethane or mixtures thereof.

Most preferably the propellant gas is 1,1,1,2-tetrafluoroethane.

The term "metered dose inhaler" or MDI means a unit comprising a canister, a secured cap covering the canister and a formulation metering valve situated in the cap. MDI system includes a suitable channelling device. Suitable channelling devices comprise, for example, a valve actuator and a cylindrical or cone-like passage through which medicament may be delivered from the filled canister via the metering valve to the nose or mouth of a patient e.g. a mouthpiece actuator.

MDI canisters generally comprise a container capable of withstanding the vapour pressure of the propellant used such as a plastic or plastics-coated glass bottle or preferably a metal canister, for example, of aluminium or an alloy thereof which may optionally be anodised, lacquer-coated and/or plastic-coated (e.g. incorporated herein by reference WO96/32150 wherein part or all of the internal surfaces of the can are coated with one or more fluorocarbon polymers optionally in combination with one or more non-fluorocarbon polymers).

The cap may be secured onto the canister via welding such as ultrasonic welding or laser welding, screw fitting or crimping. MDIs taught herein may be prepared by methods of the art (e.g., see Byron, above and WO/96/32150). Preferably the canister is fitted with a cap assembly, wherein a formulation metering valve is situated in the cap, and said cap is crimped in place.

A further aspect of the invention is a sealed container as described above capable of withstanding the pressure required to maintain the propellant as a liquid, such as a metered dose inhaler, containing therein an aerosol formulation as described above.

The formulations of the invention may be prepared by dispersal of the medicament in the selected propellant in an appropriate container, for example, with the aid of sonication or a high-shear mixer. The process is desirably carried out under controlled humidity conditions.

The chemical and physical stability and the pharmaceutical acceptability of the aerosol formulations according to the invention may be determined by techniques well known to those skilled in the art. Thus the chemical stability of the components may be determined by HPLC assay, for example, after prolonged storage of the product. Physical stability data may be gained from other conventional analytical techniques such as by leak testing, by valve delivery assay (average shot weights per actuation),

by dose reproducibility assay (active ingredient per actuation) and spray distribution analysis.

The suspension stability of the aerosol formulations according to the invention may be measured by conventional techniques, for example, by measuring flocculation size distribution using a back light scattering instrument or by measuring particle size distribution by cascade impaction or by the "twin impinger" analytical process. As used herein reference to the "twin impinger" assay means "Determination of the deposition of the emitted dose in pressurised inhalations using apparatus A" as defined in British Pharmacopaeia 1988, pages A204-207, Appendix XVII C. Such techniques enable the "respirable fraction" of the aerosol formulations to be calculated. One method used to calculate the "respirable fraction" is by reference to "fine particle fraction" which is the amount of active ingredient collected in the lower impingement chamber per actuation expressed as a percentage of the total amount of active ingredient delivered per actuation using the twin impinger method described above.

Conventional bulk manufacturing methods and machinery well known to those skilled in the art of pharmaceutical aerosol manufacture may be employed for the preparation of large scale batches for the commercial production of filled canisters. Thus, for example, in one bulk manufacturing method a metering valve is crimped onto an aluminium can to form an empty canister. The particulate medicament is added to a charge vessel and liquefied propellant is pressure filled through the charge vessel into a manufacturing vessel, together with liquefied propellant containing the surfactant. The drug suspension is mixed before recirculation to a filling machine and an aliquot of the drug suspension is then filled through the metering valve into the canister.

In an alternative process, an aliquot of the liquefied formulation is added to an open canister under conditions which are sufficiently cold such that the formulation does not vaporise, and then a metering valve crimped onto the canister.

Typically, in batches prepared for pharmaceutical use, each filled canister is checkweighed, coded with a batch number and packed into a tray for storage before release testing.

Each filled canister is conveniently fitted into a suitable channelling device, prior to use, to form a metered dose inhaler system for administration of the medicament into the lungs or nasal cavity of a patient. Metered dose inhalers are designed to deliver a fixed unit dosage of medicament per actuation or "puff", for example, in the range of 10 to 5000 micrograms of medicament per puff.

5

10

15

20

20

25

30

Administration of medicament may be indicated for the treatment of mild, moderate, severe acute or chronic symptoms or for prophylactic treatment. It will be appreciated that the precise dose administered will depend on the age and condition of the patient, the particular particulate medicament used and the frequency of administration and will ultimately be at the discretion of the attendant physician. When combinations of medicaments are employed the dose of each component of the combination will in general be that employed for each component when used alone. Typically, administration may be one or more times, for example, from 1 to 8 times per day, giving for example 1, 2, 3 or 4 puffs each time.

Suitable daily doses, may be, for example in the range 50 to 200 micrograms of salmeterol, depending on the severity of the disease.

Thus, for example, each valve actuation may deliver 25 micrograms of salmeterol (as xinafoate). Typically each filled canister for use in a metered dose inhaler system contains 60, 100, 120, 160 or 240 metered doses or puffs of medicament.

An appropriate dosing regime for other medicaments will be known or readily available to persons skilled in the art.

A further aspect of the invention provides a method of reducing drug deposition, in a pharmaceutical aerosol formulation consisting essentially of particulate medicament, e.g salmeterol xinafoate optionally in combination with another drug useful in inhalation therapy, and a liquefied propellant which is 1,1,1,2,3,3,3-heptafluoro-n-propane, 1,1,1,2-tetrafluoroethane or mixtures thereof or mixtures thereof, on valve components, especially in a metering chamber and/or sealing gaskets for use in a MDI, comprising use of at least one sealing gasket substantially constructed from a polymer of EPDM and presenting a substantially fluorinated metering chamber surface to the pharmaceutical hydrofluorocarbon aerosol formulation contained therein.

A further aspect of the invention is use of EPDM polymer in the preparation of a sealing gasket which when used in conjunction with a valve with a substantially fluorinated metering chamber surface and pharmaceutical aerosol formulation consists of or consisting essentially of particulate medicament, e.g. of salmeterol xinafoate, and a liquid propellant which is 1,1,1,2,3,3,3-heptafluoro-n-propane, 1,1,1,2-tetrafluoroethane or mixtures thereof provides the advantages described above.

The invention thus provides an MDI comprising a container, a described above, fitted with a suitable channelling device.

The use of an MDI as described above in inhalation therapy, for the treatment or prophylaxis of respiratory disorders is an alternative aspect of the invention. Specifically

10

15

20

25

the MDI system, as described above, may be used in the treatment or prophylaxis of asthma or COPD.

Furthermore the invention includes a method of treating respiratory disorders such as asthma or COPD which comprises use of an MDI as described above by a patient.

Furthermore a package comprising an MDI as described above within a flexible wrapper, said wrapper composed of a material which is substantially permeable to evacuation of propellant gas and substantially impermeable to intrusion of atmospheric moisture e.g. as described in USP 6, 119, 853 is another aspect of the invention.

Preferably the package will also contain within it a desiccant material. The desiccant material may be inside the MDI system and/or outside the MDI system.

In a further aspect the invention provides a container suitable for containing a pharmaceutical aerosol formulation comprising a canister sealed with a metering valve, said valve comprising a metering chamber having an upper and a lower sealing gasket and a valve stem, wherein the valve is sealed to the canister by means of a neck sealing gasket, characterised in that at least one gasket is substantially constructed from a polymer of EPDM and the metering chamber surface presents a substantially fluorinated surface to the formulation.

Especially preferred is a container as described above wherein the metering valve comprises a metering chamber (4) defined by walls and an upper (12) and a lower (9) sealing gasket through which pass a valve stem (7 and 8) characterised in that said two sealing gaskets are substantially constructed from a polymer of EPDM and the metering chamber surface presents a substantially fluorinated surface to a formulation containable therein.

Also especially preferred is a container as described above wherein the valve is sealed to the canister by means of a can sealing gasket (3) which is substantially constructed from EPDM polymer optionally wherein the lower (9) sealing gasket is also substantially constructed from EPDM polymer.

Most preferably all the sealing gaskets in the said metering valve are substantially constructed from EPDM polymer.

Furthermore usually the surface of surface of the metering chamber will be treated so as to present a substantially fluorinated surface to the formulation.

Thus the invention encompasses a container suitable for containing a pharmaceutical aerosol formulation comprising a canister sealed with a metering valve, said valve comprising a metering chamber having an upper and a lower sealing gasket and a valve stem characterised in that (i) the valve is sealed to the canister by means of a neck

15

20

25

30

35

sealing gasket substantially constructed from a polymer of EPDM; (ii) said upper and lower metering chamber sealing gaskets are substantially constructed from a polymer of EPDM and (iii) the metering chamber is surface treated so as to present a substantially fluorinated surface to the formulation.

Additionally the invention provides a metering valve, and use thereof, suitable for dispensing a pharmaceutical aerosol formulation comprising a metering chamber having an upper and a lower sealing gasket and a valve stem, characterised in that at least one gasket is substantially constructed from a polymer of EPDM and the metering chamber surface presents a substantially fluorinated surface to the formulation.

An exemplary valve of use according to the invention is shown in Figure 1 and comprises a valve body 1 sealed in a ferrule 2 by means of crimping, the ferrule itself being set on the neck of a container (not shown) with interposition of a gasket 3 (can seal) in a well-known manner.

The valve body 1 is formed at its lower part with a metering chamber 4, and its upper part with a sampling chamber 5 which also acts as a housing for a return spring 6. The metering chamber is constructed from a fluorinated polymer at least in part and/or a fluorinated coating according to the invention. The words "upper" and "lower" are used for the container when it is in a use orientation with the neck of the container and valve at the lower end of the container which corresponds to the orientation of the valve as shown in Figure 1. Inside the valve body 1 is disposed a valve stem 7, a part 8 of which extends outside the valve through lower stem seal 9 and ferrule 2. The stem part 8 is formed with an inner axial or longitudinal canal 10 opening at the outer end of the stem and in communication with a radial passage 11.

The upper portion of stem 7 has a diameter such that it can slide through an opening in an upper stem seal 12 and will engage the periphery of that opening sufficiently to provide a seal. Upper stem seal 12 is held in position against a step 13 formed in the valve body 1 between the said lower and upper parts by a sleeve 14 which defines the metering chamber 4 between lower stem seal 9 and upper stem seal 12. The valve stem 7 has a passage 15 which, when the stem is in the inoperative position shown, provides a communication between the metering chamber 4 and sampling chamber 5, which itself communicates with the interior of the container via orifice 26 formed in the side of the valve body 1.

Valve stem 7 is biased downwardly to the inoperative position by return spring 6 and is provided with a shoulder 17 which abuts against lower stem seal 9. In the inoperative position as shown in Figure 1 shoulder 17 abuts against lower stem seal 9 and radial

10

15.

20

25

30

passage 11 opens below lower stem seal 9 so that the metering chamber 4 is isolated from canal 10 and suspension inside cannot escape.

A ring 18 having a "U" shaped cross section extending in a radial direction is disposed around the valve body below orifice 26 so as to form a trough 19 around the valve body. As seen in Figure 1 the ring is formed as a separate component having an inner annular contacting rim of a diameter suitable to provide a friction fit over the upper part of valve body 1, the ring seating against step 13 below the orifice 26. However, the ring 18 may alternatively be formed as an integrally moulded part of valve body 1.

To use the device the container is first shaken to homogenise the suspension within the container. The user then depresses the valve stem 7 against the force of the spring 6. When the valve stem is depressed both ends of the passage 15 come to lie on the side of upper stem seal 12 remote from the metering chamber 4. Thus a dose is metered within the fluorinated metering chamber. Continued depression of the valve stem will move the radial passage 11 into the metering chamber 4 while the upper stem seal 12 seals against the valve stem body. Thus, the metered dose can exit through the radial passage 11 and the outlet canal 10.

Releasing the valve stem causes it to return to the illustrated position under the force of the spring 6. The passage 15 then once again provides communication between the metering chamber 4 and sampling chamber 6. Accordingly, at this stage liquid passes under pressure from the container through orifice 26, through the passage 15 and thence into the metering chamber 4 to fill it.

Figure 2 shows a different view of a valve in which the gasket seal and lower and upper stem seals are labelled 3, 9 and 12 respectively.

The invention will now be described further with reference the following Example which serve to illustrate the invention but is not intended to be limiting.

Example

Sample Preparation

The MDIs for which data are presented in Tables 1 to 5 were prepared in aluminium canisters coated with a PTFE/PES polymer blend as described in WO96/32150 and sealed with a Bespak valve wherein all the gaskets were made from conventional nitrile rubber (as comparator) or EPDM polymer (according to the invention) and wherein the metering chamber is composed of PBT and was conventional or surface treated with a plasma coating of a C_{1-10} perfluoroalkane.

Furthermore the said aluminium canisters contained a pharmaceutical aerosol formulation comprising 4.2mg of salmeterol xinafoate and 12g of HFA 134a.

10

15

20

25

Each device was stored at 40°C and 75% relative humidity unless otherwise stated.

Method for Determining Total Drug Content (TDC) in MDIs containing Salmeterol Xinafoate and HFA 134a

Each MDI canister tested (before use) was cooled in a freezing mixture of dry ice and methanol for approximately 5 minutes, after which it was clamped and the valve assembly removed with a suitable tube cutter. The contents of the can was quantitatively transferred into a receptacle(s) of known volume and the can, valve and valve components quantitatively washed. The combined can contents and associated washings were then assayed by HPLC and the TDC calculated. TDC values which are lower than predicted imply absorption of drug into valve components.

Mean can content is the weight of formulation contained in the canister calculated by mass difference.

Method for Determining Dose and FPM

Each MDI canister tested was put into a clean actuator and primed by firing 4 shots. Then 10 shots were fired into an Andersen Cascade Impactor which was quantitatively washed and the amount of drug deposited thereon quantified by HPLC analysis of the washings.

From this the dose delivered (the sum of the amount of drug deposited on the cascade impactor) and the FPM (the sum of drug deposited on stages two 3, 4 and 5) data were calculated. Values of FPM which are lower than expected imply one or more of the following: (i) absorption, (ii) deposition or (iii) particle growth.

The mean dose delivered is the mean of 3 dose delivered determinations. The %FPM ex-device is a measure of the dose available to the patient.

On visual inspection it was observed that the drug substance obtained from the conventional MDIs stored at 40°C 20% RH (i.e. with nitrile gaskets and a normal metering chamber [as shown in Table 1]) had the same appearance and appeared unchanged from the initial timepoint. However the drug substance from conventional MDIs stores at 40°C 75% RH was distinctly crystalline in appearance indicating some dissolution and recrystallisation.

Table 1 shows that TDC values obtained for MDIs obtained for conventional MDIs stored at stored at 40°C 75% RH and 40°C 20% RH. The former had a significantly lower TDC value than the initial timepoint and those stored under low humidity conditions.

Table 2 shows the dose delivered by the conventional MDI (control) is reduced on storage at 40°C 75% RH. The trend is very evident by the 6/7 month timepoint. The

10

15

25

30

trend is not observed in MDIs wherein all the gaskets are prepared from EPDM polymer. The trend may be present in the MDIs with a plasma treated metering chamber, however if present the trend seems to be not so pronounced as for the conventional MDI.

The FPM data for the conventional MDI employing nitrile gaskets shows a significant decrease after storage at 40°C 75% RH. This trend is reduced noticeably in addition to the initial timepoint value being higher in the MDI where all the gaskets are prepare from EPDM polymer. The data for the MDI with a plasma treated metering chamber seems to indicate an initial value for FPM higher than both the control and the EPDM polymer MDIs. However the data suggests that this value is reduced on storage at 40°C 75% RH albeit by not as much as the reduction observed for the conventional MDI.

The data in Table 3 supports the trends observed in Table 2.

The data in Table 4 shows that MDIs with gaskets substantially constructed from a polymer of EPDM and having a metering chamber with a substantially fluorinated surface provide an increase, in μg , in the dose delivered and practically eliminate the fall in dose delivered, observed on storage of the product especially under high humidity conditions, whilst simultaneously minimising the reduction in FPM observed, in comparison to conventional MDIs or those with either gaskets substantially constructed from a polymer of EPDM or having a metering chamber with a fluorinated surface.

The data in Table 5 supports the trends observed in Table 4.

The throat piece used in the Andersen Cascade Impactor to generate data contained in Table 4 and Table 5 was of the USP type. Therefore although the data was obtained using the same procedure as described above for Table 2 and Table 3 it is not directly comparable to the latter, wherein a throat manufactured for Glaxo Wellcome was used.

From the Tables it can be concluded that use of EPDM gaskets and metering chambers with a substantially fluorinated surface in MDIs containing a pharmaceutical aerosol formulation of particulate medicament, especially salmeterol xinafoate in a liquefied HFA propellant results in a formulation with improved stability in comparison to either conventional MDIs or those containing EDPM gaskets or metering chambers plasma coated with a fluorinated coating.

TABLE 1 EFFECT OF STORAGE CONDITION ON TOTAL DRUG CONTENT OF SALMETEROL INHALER

Metering	RUBBER	Storage Time	Storage	Mean TDC	Mean Can
Chamber	TYPE	Months	Condition	(mg)	Content (g)
Normal	Nitrile	10	40°C/75%RH	3.6	11.5
Normal	Nitrile	10	40°/20%RH	4.1	11.5

NOTES: All results are the mean of three individual results and TDC at initial timepoint around 4.2mg

EFFECT OF STORAGE CONDITION ON DOSE DELIVERED AND FPM OF SALMETEROL INHALER

		0	Dose Delivered (μg)	ng)		FPM (ua)	
Metering	Rubber	Initial	6 weeks	6/7 Months	Initial	6 weeks	6/7 Months
Chamber	Type						
Normal	Nitrile	18.5	16.8	13.4	9.3	7.2	5.0
Normal	EPDM	19.8	18.7	20.1	11.2	10.7	10.5
Plasma	Nitrile	21.6	•	19.7	13.0		10.3
Treated		÷					

TABLE 2

EFFECT OF STORAGE CONDITION ON MEAN DOSE DELIVERED & RANGE OF DOSE DELIVERED FOR SALMETEROL INHALERS

		Mean	Mean Dose Delivered (μg)	(bп) р	Range c	Range of Dose Delivered (µg)	ed (μg)
Metering	Rubber	Initial	6 weeks	6/7 Months	Initial	6 weeks	6/7 Months
Chamber	Туре						
Normal	Nitrile	19.1	16.8	14.5	17.1-20.7	15.4-19.2	12.8-16.1
Normal	EPDM	19.0	19.1	18.9	17.0-19.7	17.8-20.1	18.1-19.6
Plasma	Nitrile	21.2	21.0	18.4	19.9-22.1	20.5-21.5	18.0-19.4
Treated							

TABLE 3

TABLE 4 EFFECT OF STORAGE CONDITION ON DOSE DELIVERED AND FPM OF SALMETEROL INHALER

		Do	Dose Delivered (μg)	(brl) pa		FPM (μg)	a)	E.	FPM as a % of ex device Dose	of ex
Metering	Rubber	Initial	Months	ths	Initial	Mor	Months	Initial	Months	ths
Chamber	Type		င	9		က	9		m	ဖ
Normal	Nitrile	17.5	15.3	14.7	8.3	6.3	5.8	47	41	39
Normal	ЕРОМ	18.5	17.9	17.7	හ. ව.	8.4	8.0	50	47	45
Plasma Treated	Nitrile	19.9	19.0	16.8	11.1	8.4	6.5	56	44	39
Plasma Treated	EPDM	20.3	20.9	20.2	10.9	11.0	6.6	54	53	49
Treated										

EFFECT OF STORAGE CONDITION ON MEAN DOSE DELIVERED & RANGE OF DOSE DELIVERED FOR SALMETEROL INHALERS

		Me	Mean Dose Delivered (μg)	red (μg)	Ran	Range of Dose Delivered (μg)	rered (μg)
Metering	Rubber	Initial	က	9	Initial	က	9
Chamber	Туре		Months	Months		Months	Months
Normal	Nitrile	18.1	16.0	14.6	17.3-19.2	14.9–18.3	13.4-16.4
Normal	EPDM	18.3	18.7	17.4	16.8-19.9	17.7-19.5	16.4-19.2
Plasma Treated	Nitrile	19.7	19.1	17.2	19.2-20.7	18.2-19.7	16.4-18.1
Plasma Treated	EPDM	20.3	20.3	20.4	19.4-21.2	19.9-21.1	19.4-21.0

TABLE 5

Claims

10

15

- 1. A container comprising a canister sealed with a metering valve, having a metering chamber, which contains a pharmaceutical aerosol formulation consisting essentially of
- 5 (A) particulate salmeterol xinafoate optionally in combination with another drug useful in inhalation therapy, suspended in
 - (B) a liquefied propellant gas comprising 1,1,1,2,3,3,3-heptafluoro-n-propane, 1,1,1,2-tetrafluoroethane or a mixture thereof;
 - wherein the formulation is substantially free of surfactant and components having polarity higher than the liquefied propellant gas;
 - said valve characterised in that it contains one or more sealing gaskets substantially constructed from of a polymer of EPDM and the metering chamber surface presents a substantially fluorinated surface to the formulation.
 - 2. A container as claimed in claim 1 wherein the liquefied propellant gas is 1,1,1,2-tetrafluoroethane.
 - 3. A container as claimed in claim 1 or claim 2 wherein the another drug useful in inhalation therapy is fluticasone propionate or ipratropium bromide.
 - 4. A container as claimed in claim 1 or claim 2 wherein salmeterol xinafoate is the only medicament.
- 5. A container according to any one of claims 1 to 4 wherein the valve is sealed to the canister by means of a neck sealing gasket which is substantially constructed from a polymer of EPDM.
 - 6. A container according to any one of claims 1 to 5 wherein the metering valve includes a metering chamber having an upper and a lower sealing gasket and a valve stem characterised in that said two sealing gaskets are substantially constructed from a polymer of EPDM.
 - 7. A container according to any one of claims 1 to 6 wherein the metering chamber is constructed from a plastics material.
- 8. A container according to claim 7 wherein the plastics material is nylon, PBT or acetal.
 - 9. A container according to any one of claims 1 to 8 wherein the metering chamber is surface treated so as to present a substantially fluorinated surface to the formulation.
- 10. A container according to claim 9 wherein the surface treatment comprises a process of plasma coating with a C₁₋₁₀perfluoroalkane.

- A container according to claim 7 wherein the metering chamber is constructed 11. from a material selected from the group consisiting of a polyethylenetetrafluoroethylene. a polyvinyldienefluoride, a polyperfluoroalkoxyalkane, a polychlorotrifluoroethylene, a fluorinated polyethylene propylene, a copolymer of a polytetrafluoroethylene and a polyperfluoroalkoxyalkane, a copolymer of a polytetrafluoroethylene and polyhexafluoropropylene, copolymer а of a polyvinyldienefluoride and а polyhexafluoropropylene, а copolymer polytetrafluoroethylene of а and polyperfluoro(propyl vinyl ether); a blend of a polytetrafluoroethylene, а polyhexafluoropropylene a polyvinylidene fluoride, blends thereof and combinations thereof.
- 12. A container according to any one of claims 1 to 6 wherein the metering chamber is composed of a metallic material.
- 13. A container according to claim 12 wherein the metallic material is aluminium or stainless steel.
- 15 14. A container according to claim 12 or 13 wherein the metering chamber is surface treated so as to present a substantially fluorinated surface to the formulation.
 - 15. A container according to claim 14 wherein the surface treatment comprises a process of applying a coating of a fluorocarbon polymer optionally in combination with a non-fluorocarbon polymer.
- 20 16. A container according to claim 14 or claim 15 wherein fluorocarbon polymer is selected from FEP and PTFE.
 - 17. A container according to any one of claims 14 to 16 wherein the coating is a coating of FEP.
- 18. A container according to any one of claims 14 to 16 wherein the coating is a coating of a blend of PTFE and PES.
 - 19. A container according to any one of claims 1 to 18 wherein the canister is composed of aluminium.
 - 20. A container according to claim 19 wherein the canister is surface treated so as to present a substantially fluorinated surface to the formulation.
- 30 21. A container according to claim 20 wherein the canister is surface treated by coating with a fluorocarbon polymer optionally in combination with a non-fluorocarbon polymer.
 - 22. A container according to claim 20 or claim 21 wherein fluorocarbon polymer is selected from FEP and PTFE.

- 23. A container according to any one of claims 20 to 22 wherein the coating is a coating of FEP.
- 24. A container according to any one of claims 20 to 22 wherein the coating is a coating of a blend of PTFE and PES.
- 5 25. A metered dose inhaler comprising a container according to any one of claims 1 to 24 fitted with a suitable channelling device.
 - A method of treatment of asthma or COPD which comprises use of a metered dose inhaler according to claim 25 by a patient.
- 27. A package comprising a metered dose inhaler according to claim 25 contained within a flexible wrapper said wrapper composed of a material which is substantially permeable to evacuation of propellant gas and substantially impermeable to intrusion of atmospheric moisture.
 - 28. A package according to claim 27 characterised in the flexible wrapper also contains within it a desiccant material.
- 15 29. A package according to claim 28 characterised in the can contains within it a desiccant material.
 - 30. A container suitable for containing a pharmaceutical aerosol formulation comprising a canister sealed with a metering valve, said valve comprising a metering chamber having an upper and a lower sealing gasket and a valve stem, wherein the valve is sealed to the canister by means of a neck sealing gasket, characterised in that at least one gasket is substantially constructed from a polymer of EPDM and the metering chamber surface presents a substantially fluorinated surface to the formulation.
 - 31. A container suitable for containing a pharmaceutical aerosol formulation according to claim 30 wherein the upper, lower and neck sealing gaskets are substantially constructed from a polymer of EPDM.
 - 32. A container suitable for containing a pharmaceutical aerosol formulation according to claim 30 or claim 31 wherein the metering chamber is surface treated so as to present a substantially fluorinated surface to the formulation.
 - 33. A container according to claim 32 wherein the surface treatment comprises a process of plasma coating with a C₁₋₁₀perfluoroalkane.
 - 34. A container according to any one of claims 30 to 33 wherein the metering chamber is constructed from a plastics material.
 - 35. A container according to claim 34 wherein the plastics material is nylon, PBT or acetal.

25

- 36. A container according to any one of claims 30 to 35 wherein the canister is composed of aluminium.
- 37. A container according to claim 36 wherein the canister is surface treated so as to present a substantially fluorinated surface to the formulation.
- 5 38. A container according to claim 37 wherein the canister is surface treated by coating with a fluorocarbon polymer optionally in combination with a non-fluorocarbon polymer.
 - 39. A container according to any one of claims 30 to 38 which contains a pharmaceutical aerosol formulation comprising a particulate medicament and a liquefied propellant gas of 1,1,1,2,3,3,3-heptafluoro-n-propane, 1,1,1,2-tetrafluoroethane or mixtures thereof.
 - 40. A container according to claim 39 wherein the propellant gas is 1,1,1,2-tetrafluoroethane.
- 41. A container according to claim 39 or claim 40 wherein the particulate medicament is selected from salmeterol xinafoate, fluticasone propionate, albuterol or a salt thereof, beclomethasone dipropionate, formoterol or a salt thereof, ipratropium bromide, budesonide, sodium cromoglycate and combinations thereof.
 - 42. A container according to claim 41 wherein the particulate medicament is salmeterol xinafoate optionally in combination with fluticasone propionate.

1/2

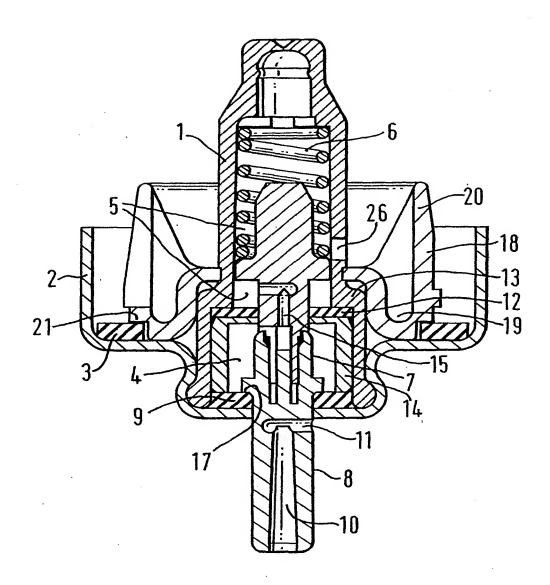


FIG. 1.

SUBSTITUTE SHEET (RULE 26)

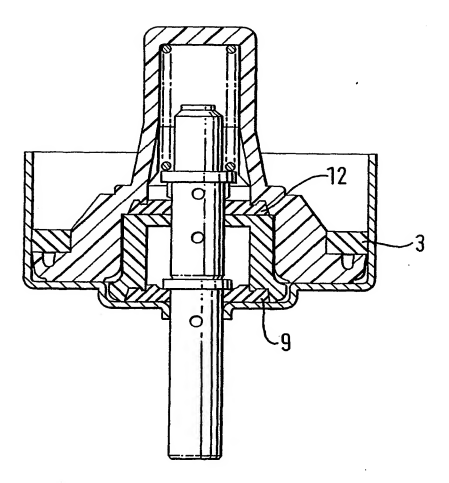


FIG. 2.

SUBSTITUTE SHEET (RULE 26)

PCT/GB 01/05749

		FCI/UD	01/05/49
A. CLASSI IPC 7	IFICATION OF SUBJECT MATTER A61M15/00 A61K9/00 B65D83/	14	
According to	o International Patent Classification (IPC) or to both national classific	cation and IPC	<u></u>
B. FIELDS	SEARCHED	~	
Minimum do IPC 7	ocumentation searched (classification system followed by classificat $A61M-A61K-B65D$	ion symbols)	
	tion searched other than minimum documentation to the extent that state of the extent that state		
	ternal, WPI Data, PAJ	ise and, where practical, segion terms (usea)
C. DOCUME	ENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the rel	levant passages	Relevant to claim No.
Х	WO 95 02651 A (MINNESOTA MINING 8		1,2,4-6,
Υ	26 January 1995 (1995-01-26) page 11, line 19 - line 27; claim	ns 15,19	19,25 3,7-18, 20-27, 30-40
Y,P	WO 01 76601 A (RICHARDS DAVID HUG GROUP LTD (GB); JENKINS RICHARD 3 18 October 2001 (2001-10-18) abstract	ЭН ;GLAXO JOHN (U)	3
Y	WO 99 47195 A (GLAXO GROUP LTD ;F MICHAEL THOMAS (US); SCHULZE MARK () 23 September 1999 (1999-09-23) claims	7-11, 30-40	
		-/	
X Furth	er documents are listed in the continuation of box C.	X Patent family members are lis	sled in annex.
° Special cat	egories of cited documents :	*T* later document published after the	international filing date
conside	nt defining the general state of the art which is not ered to be of particular relevance ocument but published on or after the international	or priority date and not in conflict value of the understand the principle of invention "X" document of particular relevance; the	with the application but r theory underlying the the claimed invention
"L" documer	ate which may throw doubts on priority claim(s) or	cannot be considered novel or can involve an inventive step when the	nnot be considered to e document is taken alone
citation "O" docume	or other special reason (as specified) ont referring to an oral disclosure, use, exhibition or	"Y" document of particular relevance; it cannot be considered to involve a document is combined with one or	n inventive step when the r more other such docu-
"P" document later the	nt published prior to the international filling date but	ments, such combination being ob in the art. *& document member of the same pate	·
	actual completion of the international search	Date of mailing of the international	
19	9 April 2002	03/05/2002	
Name and m	nailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2	Authorized officer	
	NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Villeneuve, J-M	

Form PCT/ISA/210 (second sheet) (July 1992)

PCT/GB 01/05749

0.10	ALL DECOMPTANTS CONCURRENTS TO BE THE TANK	1C1/0B 01/03/49
	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	Dolor-st to -1-1- to
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Υ	WO 96 32150 A (GLAXO WELLCOME INC ;ASHURST IAN C (US); HERMAN CRAIG S (US); LI LI) 17 October 1996 (1996-10-17) cited in the application page 6, line 30 -page 7, line 20; claims	12-18, 20-24
Υ	WO 00 37336 A (GARRILL KARL ANDREW ;GLAXO GROUP LTD (GB); WALKER RICHARD IAN (GB)) 29 June 2000 (2000-06-29) cited in the application claims 1,2,9,10	25–27
А	WO DO 56632 A (ANDERSON GREGOR JOHN MCLENNAN; GODFREY JAMES WILLIAM (GB); GLAXO G) 28 September 2000 (2000-09-28) page 9, last paragraph; claims 25-27,33,34; figures	3,30-40
A	EP 0 990 437 A (GLAXO GROUP LTD) 5 April 2000 (2000-04-05) paragraphs '0005!,'0006!,'0009!,'0012!,'0015!	1
A	US 6 089 256 A (WARBY RICHARD JOHN) 18 July 2000 (2000-07-18) claims	12-18

Form PCT/ISA/210 (continuation of second sheet) (July 1992)

PCT/GB 01/05749

				101740	01/05/49
Patent document cited in search report		Publication date		Patent family member(s)	Publication date
WO 9502651	A	26-01-1995	AU DE DE DE EP ES JP WO US	680530 B2 7395694 A 9422364 U1 69412626 D1 69412626 T2 0708805 A1 2119219 T3 3172190 B2 9500300 T 9502651 A1 5836299 A	31-07-1997 13-02-1995 14-09-2000 24-09-1998 28-01-1999 01-05-1996 01-10-1998 04-06-2001 14-01-1997 26-01-1995 17-11-1998
WO 0176601	A	18-10-2001	AU WO	7391901 A 0176601 A2	23-10-2001 18-10-2001
WO 9947195	А	23-09-1999	AU BR CA CN EE WO EP HR JP NO PL SK TR	3145799 A 9908766 A 2324524 A1 1293580 T 200000549 A 9947195 A1 1064040 A1 20000610 A1 0101223 A2 2002506696 T 20004642 A 342972 A1 13742000 A3 200002663 T2	11-10-1999 14-11-2000 23-09-1999 02-05-2001 15-02-2002 23-09-1999 03-01-2001 30-04-2001 28-09-2001 05-03-2002 18-09-2000 16-07-2001 06-08-2001 21-12-2000
WO 9632150	A	17-10-1996	AP AU BG BR CCN CZ EE HU JP NO NZ PL SK TR WO US	979 A 718263 B2 5481196 A 102022 A 9604977 A 2217954 A1 1186447 A 9703260 A3 9700374 A 0820323 A1 9802391 A2 11509434 T 974736 A 306280 A 322771 A1 138997 A3 9701169 T1 9632150 A1 6143277 A	28-06-2001 13-04-2000 30-10-1996 31-07-1998 09-06-1998 17-10-1996 01-07-1998 18-02-1998 15-06-1998 28-01-1998 01-02-1999 24-08-1999 11-12-1997 29-07-1999 16-02-1998 08-04-1998 21-03-1998 17-10-1996 07-11-2000
WO 0037336	A	29-06-2000	US US AU BR CN EP NO TR	6119853 A 6179118 B1 1830300 A 9916336 A 1334774 T 1150905 A1 20012997 A 200101959 T2	19-09-2000 30-01-2001 12-07-2000 11-09-2001 06-02-2002 07-11-2001 15-08-2001 21-01-2002

Form PCT/ISA/210 (patent family annex) (July 1992)

PCT/GB 01/05749

				101/4	5 U1/U5/49
Patent documer cited in search rep		Publication date		Patent family member(s)	Publication date
WO 0037336	. А	· · · · · ·	WO US US	0037336 A1 6352152 B1 6315112 B1	29-06-2000 05-03-2002 13-11-2001
WO 0056632	• А	28-09-2000	AU BR CN WO	3656200 A 0009227 A 1344217 T 0056632 A2	09-10-2000 26-12-2001 10-04-2002 28-09-2000
			EP	1144272 A2	17-10-2001
EP 0990437	• А	05-04-2000	EP EP	1066828 A1 0990437 A1	10-01-2001 05-04-2000
			EP	0756868 A2	05-02-1997
			ΑU	663904 B2	26-10-1995
ľ			ΑU	3085092 A	19-07-1993
			BG	62119 B1	31-03-1999
			BG	98803 A	28-02-1995
			DE	69224656 D1	09-04-1998
•			DE	69224656 T2	23-07-1998
j			DK	616523 T3	28-09-1998
			EP HK	0616523 A1 1004711 A1	28-09-1994 28-07-2000
			JP	3026840 B2	27-03-2000
			JP	7502033 T	02-03-1995
			NO	942185 A	10-06-1994
			NO	20001227 A	10-06-1994
			RU	2129424 C1	27-04-1999
			SK	67494 A3	08-03-1995
			US	6238647 B1	29-05-2001
			US	5674472 A	07-10-1997
			US	6303103 B1	16-10-2001
	•		US	6251368 B1	26-06-2001
			AP	402 A	22-08-1995
			AT	163539 T	15-03-1998
			AT	201587 T	15-06-2001
]			BG	102689 A	26-02-1999
			CA	2125667 A1	24-06-1993
			CA	2303685 A1	24-06-1993
			CZ	9401430 A3	15-03-1995 05-07-2001
1			DE DE	69231857 D1 69231857 T2	29-11-2001
			DK	756868 T3	10-09-2001
			WO	9311743 A1	24-06-1993
1			ËS	2113444 T3	01-05-1998
			ES	2158988 T3	16-09-2001
1			HU	67534 A2	28-04-1995
			HU	9500331 A3	28-09-1995
			IL	104068 A	30-10-1998
			JP	11310533 A	09-11-1999
, Á.			MX	9207205 A1	01-11-1993
Ţ.			NZ	246044 A	26-01-1996
			OA OT	9926 A	15-09-1994
			PT	756868 T	30-11-2001
			SG	74042 A1	18-07-2000
			US	5674471 A	07-10-1997 14-10-1997
1			US US	5676929 A	14 - 10-1997
			US	5658549 A 5683676 A	19-08-1997 04-11-1997
}			uş	3003070 A	04 .11122/
Form PCT/(SA/210 (patent family and	4411				

Form PCT/ISA/210 (patent family annex) (July 1992)

PCT/GB 01/05749

Patent document cited in search report		Publication date		Patent family member(s)	Publication date
EP 0990437	Α		US	5653962 A	05-08-1997
US 6089256	A	18-07-2000	DE FR FR GB GB US	19835273 A1 2767801 A1 2779705 A1 2328932 A ,B 2338951 A ,B 6095182 A	04-03-1999 05-03-1999 17-12-1999 10-03-1999 12-01-2000 01-08-2000

Form PCT/ISA/210 (patent family annex) (July 1992)

Draft - Not for Implementation

Guidance for Industry

Metered Dose Inhaler (MDI) and Dry Powder Inhaler (DPI) Drug Products

Chemistry, Manufacturing, and Controls Documentation

DRAFT GUIDANCE

This document is being distributed for comment purposes only.

Comments and suggestions regarding this draft document should be submitted within 90 days of publication in the *Federal Register* of the notice announcing the availability of the draft guidance. Submit comments to Dockets Management Branch (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. All comments should be identified with the docket number listed in the notice of availability that publishes in the *Federal Register*.

For questions regarding this draft document, contact Guirag Poochikian, Ph.D., (301) 827-1050.

U.S. Department of Health and Human Services
Food and Drug Administration
Center for Drug Evaluation and Research (CDER)
October 1998
CMC

X:\CDERGUID\2180DFT.WPD November 13, 1998